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(54) Method for the marking of neurones or neurone substructures

(57) The present invention relates to an *ex vivo* method for marking neurones or neurone substructures of a mammal, comprising detecting the expression of a protein represented by SEQ ID NO: 4 (*Mus musculus* BADLAMP) or by a homologous sequence which presents at

least 60% identity to SEQ ID NO: 4, provided that the homologous sequence represents a protein which can be found associated to plasma and/or vesicular membranes within cortical neurones of said mammal.

Description

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Field of the invention

5 [0001] The present invention relates to a method for marking neurones or neurone substructures.

Background of the invention

[0002] The separation of membrane lipids into different phases creates diverse microenvironments within a biological membrane. In particular, cholesterol is believed to condense with saturated phosphatidylcholine (PC) and sphingomyelin (SM) to form minute patches (40-100 nm wide) of lipids in a liquid-ordered phase, creating specialized lipid microdomains called "rafts" within the disordered fluid phase formed by unsaturated lipids. These ordered domains control the access and egress of subsets of membrane proteins, regulating signaling systems at the cell surface. Liquid-ordered domains resist solubilization in non-ionic detergents.

[0003] In neurons, several types of microdomains have been shown to be distinguishable by the partitioning of different membrane associated proteins such as Thy-1, the major GPI-anchored protein of mature neurons or the GPI-anchored prion protein (PrP). Thy-1 and PrP are found on the neuronal surface in different, albeit often closely adjacent domains. These differences in surface localization are reflected in the different functions and trafficking of these proteins. Thy-1 inhibits the activity of Src family kinases attached to the inner leaflet of rafts, undergoes relatively slow internalization while PrP, in concert with the neural cell adhesion molecule (NCAM), is rapidly and constitutively endocytosed on neurons and induces axonal outgrowth via the activation of Fyn Kinases. Partition in microdomains has been shown to regulate the function of many neuronal proteins such as potassium-gated channel isoforms, type-1 cannabinoid receptors (CB1R) or NCAM. Lipid microdomains and their organization play therefore a key role in neuronal development and function.

[0004] A body of evidence links lipid rafts to the generation and function of synapses in the brain. For example, gliaderived cholesterol has been shown to be important for synapse formation and lipid-rich microdomains have been found abundantly on dendrites and associated to postsynaptic components like for example AMPA receptors. BDNF exposure has been shown to induce the selective translocation of the TrkB receptor into lipid rafts of cultured cortical neurons. Finally, the ubiquitinin related system has been shown to be present in lipid rafts of postsynaptic densities and suggested to be crucially involved in their regulation.

[0005] Lysosomal Associated Membrane Proteins (LAMPs) form a family of heavily glycosylated membrane glycoproteins which are mostly localized in lysosome and have been shown to play an important role in their biogenesis (Eskelinen et al. (2003) Trends Cell Biol 13:137).

[0006] The present invention arises from the discovery that in mice brain, a previously unknown transmembrane protein, closely related to the LAMP family, is specifically expressed in cortical neurons of layers II, III and V. Furthermore, the inventors have shown that this protein is enriched in defined zones along the different neuronal projections detected both in brain section and in cortical neuron cultures by accumulating in intracellular vesicles, which do not contain any known markers of the classical neuronal intracellular transport pathways, thus defining a new class of intracellular vesicles.

Summary of the invention

[0007] As such the present invention relates to a method, preferably an *ex vivo* method, for marking neurones or neurone substructures of a mammal, comprising detecting the expression of a protein represented by SEQ ID NO: 4 (*Mus musculus* BAD-LAMP) or by a homologous sequence which presents at least 60%, preferably at least 70%, more preferably at least 80%, identity to SEQ ID NO: 4, provided that the homologous sequence represents a protein which can be found associated to plasma and/or vesicular membranes within cortical neurones of said mammal.

[0008] In an embodiment of the above-defined method, the expression of the protein is detected by a ligand comprising at least one binding moiety, which at least one binding moiety is specific for said protein.

[0009] In another embodiment of the above-defined method, the ligand further comprises at least one detectable marker.

[0010] In a further embodiment of the above-defined method, the ligand consists of at least one binding moiety linked to at least one detectable marker.

[0011] In another further embodiment of the above-defined method, the ligand consists of at least one binding moiety. [0012] In another embodiment of the above-defined method the messenger RNA (mRNA) which encodes the above-defined protein, or a precursor thereof, such as a pre-mRNA, is detected.

Description of the figures

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Figure 1 represents the peptide sequence of mouse BAD-LAMP (from the N-terminus to the C-terminus). The first underlined part of the sequence (starting from the N-terminus) represents the transmembrane domain and the second underlined part of the sequence represents a consensus internalization sequence and endosomal targeting signal.

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Figure 2 represents a northern blot of various mouse tissue samples (on top) using a radio-labelled BAD-LAMP mRNA-specific probe.

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Figure 3 represents a northern blot of various human tissue samples (on top) using a radio-labelled BAD-LAMP mRNA-specific probe.

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Figure 4 represents an immunoblot performed with a polyclonal antibody raised against a predicted peptide of the BAD-LAMP cytoplasmic tail. Lysates of mouse adult cortex and BAD-LAMP transfected HeLa cells display several bands migrating above 35 kDa, while no reactivity is observed in control untransfected cells, the lowest form accumulating in the transfected cells is probably due to ER accumulation.

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Figure 5 represents an immunoblot performed with a polyclonal antibody raised against a predicted peptide of the BAD-LAMP cytoplasmic tail. HeLa cells transfected with FLAG-BAD-LAMP cDNA are lysed and immunoprecipitated with anti-FLAG antibody. Immunoprecipitated material is treated with endoglycosidase H (endo H) or N-glycosidase F (N-gly F) prior immunoblotting with anti-BAD-LAMP. In untransfected cells (NC), just the anti-FLAG IgG band (arrow) is observed, while several isoforms of BAD-LAMP are immunoprecipitated in transfected cells (Wt). Endo H treatment shows that the major band is endo H sensitive (gH) thus probably accumulating in the ER. The higher molecular weight bands are all N-gly F sensitive (gF). All BAD-LAMP isoforms are glycosylated and its native molecular weight is as predicted close to 32kDa (g0). The anti-FLAG IgG band are also N-gly F sensitive, arrows.

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Figure 6 represents an immunoblot performed with a polyclonal antibody raised against a predicted peptide of the BAD-LAMP cytoplasmic tail. After fractionation and isolation of cortical membranes, BAD-LAMP is found present exclusively in the membrane pellet (MbP) and not in the supernatant (Sup). Control syntaxin 6 and syntaxin 13 were also found in the membrane pellet, while Rab3 was distributed in the two fractions due to its shuttling nature.

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Figure 7 represents an immunoblot performed with a polyclonal antibody raised against a predicted peptide of the BAD-LAMP cytoplasmic tail. Post-natal mouse cortex extracts were immunoblotted with BAD-LAMP polyclonal antibody. BAD-LAMP expression is increased from birth to adulthood (P0 to Ad).

Figure 8 represents *in situ* hybridization for BAD-LAMP on coronal post-natal brain sections from P2 to P12. Hemisections are presented. Bad-lamp expression signal appears at P2 in the cingulate cortex (ci) and extended ventrally during the first post-natal weeks as a superficial and a deep band in the cortex (ctx). In subcortical structure, BAD-LAMP is expressed transiently in the caudate putamen (cp). Bar: 500 µm.

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Figure 9 represents *in situ* hybridization for BAD-LAMP (top panel), Cux2 (middle) and ER81 (bottom) on adult brain coronal hemisections.

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Figure 10 represents high magnification views of *in situ* hybridization on wild type cortex shown in Figure 9 and *in situ* hybridization for BAD-LAMP in Scrambler cortex. BAD-LAMP is expressed in layers II, II and V, but excluded from layers IV and VI. In the Scrambler cortex, the entire cortex appears disorganized. However, the typical inversion of cortical layers is reflected by the altered BAD-LAMP staining, demonstrating that projection neurons express the protein.

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Figure 11 represents a list of the different molecules imaged in cortical neurones by confocal microscopy is given along with their level of co-localization with BAD-LAMP.

Figure 12 represents a subcellular Percoll gradient fractionation showing that in transfected HeLa cells BAD-LAMP is mostly localized in light density fractions and is absent from heavy density fractions containing the lysosomal enzyme β-hexosaminidase.

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Figure 13 shows a representative FACS staining experiment (of 3), showing the level of internalised FLAG antibody against time at the surface of FLAG-BAD-LAMP transfected HeLa cells. The timing of internalization and the recovery of surface staining at 10 min indicates the BAD-LAMP recycles in HeLa cells.

Figure 14 represents the percentage of amino acid identity of *Rattus norvegicus* (SEQ ID NO: 6), *Bos Taurus* (SEQ ID NO: 8), *Homo sapiens* (SEQ ID NO: 2), *Takifugu rubripes* (SEQ ID NO: 11), *Tetraodon nigroviridis* (SEQ ID NO: 12), *Gallus galius* (SEQ ID NO: 13), *Drosophila melanogaster* (SEQ ID NO: 14), and *Anopheles gambiae* (SEQ ID NO: 15) BAD-LAMPs with respect to *Mus musculus* (SEQ ID NO: 4) BAD-LAMP.

Detailed description of the invention

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[0014] The method according to the invention is liable to be implemented *in vivo* or *ex vivo*. When implemented *ex vivo* the method can be applied to tissue samples such as biopsies or histological sections. The method can also be implemented in combination with other methods for marking neurones or neurone substructures already known in the art.

[0015] As intended herein the expression "marking neurones or neurone substructures" relates to the ability to discriminate neurones or neurone substructures expressing the protein from those which do not.

[0016] Neurones which express the protein are localized within particular cerebral cortical layers of the mammal. These particular cerebral cortical layers are preferably selected from the group constituted of layers II, III and V. The nomenclature of cerebral cortical layers is well known to one skilled in the art and is notably described in Molnar et al. (2006) European Journal of Neuroscience 23:921-934. Advantageously, the method according to the invention thus provides for a simple method to detect cerebral cortical layers of a mammal, for instance on histological sections of a mammal's brain.

[0017] Neurone substructures liable to be detected according to the invention encompass particular regions of neurones, such as cellular body, neurites, growth cones or synapses, and cellular components, such as the plasma membrane, or intracellular vesicles.

[0018] The method according to the invention can be implemented for any mammal. For each mammal, the expression of a protein of said mammal having a sequence homologous to the sequence SEQ ID NO: 4 should be detected. As intended herein, a homologous sequence shares at least 60%, preferably at least 70%, more preferably at least 80%, identity with SEQ ID NO: 4 and represents a protein of the mammal which can be found associated to plasma and/or vesicular membranes within cortical neurones of said mammal. More particularly, said protein of the mammal can be found essentially co-localizing with GM-1 ganglioside upon cholesterol depletion of the neurones of said mammal. Cholesterol depletion can be implemented by treating neurones with cholesterol oxydase as described in the following Examples.

30 [0019] As intended herein a first peptide sequence which presents at least x % identity to a second peptide sequence means that x % represents the number of amino acids of the first sequence which are identical to their paired amino acids of the second sequence when the two sequences are optimally aligned pairwise, with respect to the total length of the second peptide sequence. As intended herein two sequences are said to be optimally aligned pairwise when x is maximal.

[0020] Procedures for finding if a protein is associated to detergent-insoluble lipid microdomains within neurones of said mammal are well-known to one skilled in the art and are notably illustrated in the following examples.

[0021] In particular, when the mammal is:

- a mouse, more particularly Mus musculus, the sequence of the protein is preferably SEQ ID NO: 4;
- a human, the sequence of the protein is preferably SEQ ID NO: 2;
- a rat, more particularly Rattus norvegicus, the sequence of the protein is preferably SEQ ID NO: 6;
- a bovine, more particularly *Bos taurus*, the sequence of the protein is preferably SEQ ID NO: 8;
- an ape, more particularly Pan troglodytes, the sequence of the protein is preferably SEQ ID NO: 10.

[0022] As will be apparent to one skilled in the art, a "homologous sequence" according to the invention, also encompasses natural variants of SEQ ID NO: 2, 4, 6, 8 and 10 in the respective corresponding mammalian species.

[0023] As intended herein the expression "detecting the expression of a protein" encompasses detecting the protein in itself or the messenger RNA (mRNA) which encodes said protein, or precursors thereof, such as a pre-mRNA.

[0024] When it is sought to detect the protein in itself, a ligand specific for said protein may be used.

[0025] As intended herein a "ligand" refers to any compound liable to specifically bind to the protein as defined above. The part or parts of the ligand which effectively bind to the protein as defined above are named "binding moiety" or "binding moieties". A ligand according to the invention can thus comprise or be constituted of one or several binding moieties. In particular, when a ligand according to the invention comprises one or several binding moieties, it can also comprises at least one "detectable marker", that is a moiety the presence of which can be readily evidenced according to methods well known to the man skilled in the art.

[0026] Preferably, in the ligands according to the invention, the at least one binding moiety is specific for a protein represented by a sequence selected from the group consisting of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, SEQ ID NO: 8, and SEQ ID NO: 10.

[0027] Also preferably, the at least one binding moiety is selected from the group consisting of an antibody, an antigen-specific antibody fragment, a scFv, or an aptamer.

[0028] As intended herein an "antigen-specific antibody fragment" relates to an antibody fragment which retains its specific binding properties towards an isolated protein according to the invention. Such fragments notably encompass Fab fragments (which can be produced by papain cleavage of antibodies), F(ab')₂ fragments (which can be produced by pepsin cleavage of antibodies) or Fab' fragments (which can be produced by pepsin cleavage of antibodies followed by a reducing treatment).

[0029] A "scFv" relates to a single-chain variable fragment of an antibody, that is an immunoglobulin short chain variable region and an immunoglobulin large chain variable region linked together by a peptide.

[0030] An "aptamer" relates to a nucleic acid molecule, in particular a ribonucleic acid molecule.

[0031] Methods for producing an antibody, an antigen-specific antibody fragment, a scFv, or an aptamer are well-known to the man skilled in the art.

[0032] More preferably the at least one binding moiety is a monoclonal antibody.

[0033] General methods for producing monoclonal antibodies are well known to the man skilled in the art and are notably described by Kohler & Milstein (1975) Nature 256:495 or illustrated in the following Examples.

[0034] Most preferably the monoclonal antibody is produced by one of the following hybridomas:

- a hybridoma deposited at the Collection Nationale de Cultures de Microorganismes (CNCM, Paris, France) under the Budapest Treaty on October 19, 2006 with accession number I-3679, or
- a hybridoma deposited at the Collection Nationale de Cultures de Microorganismes (CNCM, Paris, France) under the Budapest Treaty on October 19, 2006 with accession number 1-3679.

[0035] Preferably, the above-defined at least one detectable marker is selected from the group consisting of an enzyme, a chromophore, or a fluorophore.

[0036] Alternatively, to detect the expression of a protein as defined above, it may be sought detect the messenger RNA (mRNA) which encodes the above-defined protein, or a precursor thereof, such as a pre-mRNA.

[0037] When the mammal is:

- a mouse, more particularly *Mus musculus*, the mRNA, or precursor thereof, should preferably comprise the sequence SEQ ID NO: 3;
- a human, the mRNA, or precursor thereof, should preferably comprise the sequence SEQ ID NO: 1;
- a rat, more particularly *Rattus norvegicus*, the mRNA, or precursor thereof, should preferably comprise the sequence SEQ ID NO: 5;
- a bovine, more particularly *Bos Taurus*, the mRNA, or precursor thereof, should preferably comprise the sequence SEQ ID NO: 7;
- an ape, more particularly Pan troglodytes, the sequence of the protein is preferably SEQ ID NO: 9.

[0038] Several methods well-known to one skilled in the art may be used to detect the messenger RNA (mRNA) which encodes the above-defined protein or precursors thereof, such as a pre-mRNA.

[0039] In particular a detection nucleic which is liable to hybridize with a nucleic acid encoding the protein may be used.

[0040] Preferably, the detection nucleic acid hybridizes under stringent conditions with a nucleic acid encoding the protein.

[0041] Stringent conditions for nucleic acid hybridization are well known to the man skilled in the art and are notably described in Sambrook & Rusell (2001) "Molecular Cloning: A Laboratory Manual", 3rd edition, Cold Spring Harbor Laboratory Press. By way of example, suitable stringent conditions according to the invention with respect to a given nucleic acid consist in a hybridization carried out at temperatures 20°C below the calculated melting temperature (Tm) for said nucleic acid in the same medium than that used for calculating the Tm.

[0042] More preferably, the detection nucleic acid is selected from the group constituted of nucleic acids comprising or constituted of the complementary sequences of SEQ ID NO: 1, 3, 5, 7, and 9 or fragments thereof having at least 9 nucleotides.

[0043] Alternatively, the expression of the protein is detected by a nucleic acid amplification method.

[0044] Nucleic acid amplification methods suitable for implementing the method according to the invention are well-known to one skilled in the art and notably encompass reverse-transcriptase polymerase chain reaction (RT-PCR) methods.

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EXAMPLES

Example 1

5 Materials and methods

Bioinformatics

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[0045] BAD-LAMP protein sequence ID in Ensembl database: ENSMUSP00000061180 All LAMPs sequences were aligned using CLUSTALW package (EBI) and results were treated with TreeView for phylogeny. Image analysis was performed with the Image J software and the plugin JacoP designed by Fabrice Cordelières, Institut Curie, Orsay (France).

Animals and Tissues

15 [0046] All animals were treated according to protocols approved by the French Ethical Committee.

[0047] CD1 mice (Iffa-Credo, France) were used to determine BAD-LAMP expression pattern. Disabled-1 deficient Scrambler mice have been purchased from Jackson Laboratories.

[0048] The day of the vaginal plug appearance was considered as E0.5 and the day of the birth is termed postnatal (P) 0.

[0049] For in situ hybridisation and immunohistochemistry, postnatal and adult brains were collected after the animals were anaesthetized with a lethal dose of Rompun/Imalgen 500 and intracardiacally perfused with 4% paraformaldehyde (PFA). Brains were further fixed in 4% PFA overnight. Adult brains were sectioned at 80 μ m on a vibratome while P2 to P12 brains were cryoprotected in 20% sucrose/PBS, frozen in OCT compound and sectioned at 16 μ m on a cryostat. Sections collected on Superfrost slides were treated as described below.

25 Molecular biology

[0050] Northern blot was done with FirstChoice Northern Blot Mouse Blot I (Ambion) using a probe corresponding to exons 4, 5 and 6 of BAD-LAMP (clone IMAGE 2588577). 2 µg of Trizol extracted total mouse cortex RNA was used for reverse transcription with oligo(dT) primers. The cDNAs coding for BAD-LAMP were amplified after 30 cycles of PCR using Taq polymerase. Sense primer was ACC GGC CAC TTT GAG GGA (SEQ ID NO: 16) and antisense GGG GCC GCC TTT GCA GCA (SEQ ID NO: 17) (1,5 kbp). PCR products were cloned into pGEM-Teasy plasmid (Promega).

[0051] BAD-LAMP-GFP fusion construct was done in pEGFP-NI vector (Clontech). FLAG-BAD-LAMP was constructed using pTEJ-8-HAFLAG plasmid (Didier Marguet, Marseille).

[0052] Tyrosine mutant of BAD, FLAG-BAD-Tyr-276-Ala was realized by targeted PCR mutagenesis.

[0053] FLAG-BAD-LAMP cDNAs were transferred into pCX-MCS2 plasmid, a pCAAGS derived plasmid with an extended cloning site (a kind gift of Xavier Morin, Marseille). Dynamin-GFP wt plasmid and dynamin-GFP A44K were kindly given by M. McNiven, Rochester. RNAi constructs pSUPER AP-2 μ2 and pSUPER control were a gift from Dr Philippe Benaroch, Paris.

40 In situ hybridization and immunohistochemistry

[0054] IMAGE clone 2588577 was used to make an antisense RNA probe. Antisense RNA probe for BADLAMP, Cux2 (Zimmer et al., 2004) and ER81 (Lin et al., 1998) were generated using the Dig-RNA labelling kit (Roche).

[0055] Single *in situ* hybridization and combined *in situ* hybridization with immunohistochemistry were described previously (Tiveron et al., 1996) (Zimmer et al., 2004) for all probes and the NeuN monoclonal mouse IgG (MAB377, Chemicon).

Antibodies and immunocytochemistry.

[0056] A polyclonal rabbit anti-BAD-LAMP was raised in rabbit against two peptides of BADLAMP cytoplasmic tail, KMTANQVQIPRDRSQC (SEQ ID NO: 18) and KQIPRDRSQYKHMC (SEQ ID NO: 19).

[0057] Antisynaptotagmin I and anti-Rab3a/b antibodies was obtained from Dr P. Di Camilli, New Haven, anti-FLAG M2 antibody and anti-®-tubulin-Cy3 and FITC-cholera toxin B subunit (GM-1 staining) were obtained from Sigma, anti-VAMP2 was from SYSY, antisyntaxin 6 was from BD Transduction Laboratories, anti-PrP (6H4) was from Prionics, anti-syntaxin 13 was from Stressgen, anti-Thy1 was from Michel Pierres, Marseille, human Alexa 568-Tf was from molecular probes, mouse FITC-Tf was from Rockland, anti-NCAM H28 from Dr. C. Goridis, Paris, Anti-Ti-VAMP from Dr. T. Galli, Paris, Rat anti mouse LAMP2 from Dr I. Mellman, New Haven and anti human LAMP-1 from Abcam. All FITC and Cy3-5 secondary antibodies were from Jackson Immunoresearch. All Alexa secondary antibodies were from Molecular Probes

(USA).

[0058] Immunofluorescence and confocal microscopy was performed with a Zeiss LSM 510 as described (Cappello et al., 2004). Vibratome adult brain sections were immunostained with rabbit anti-BAD-LAMP and mouse anti-MAP2.

5 Cell culture

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[0059] HeLa cells were grown in DMEM containing 10% FCS.

[0060] Cortical neurons were prepared from E15.5 embryonic cortices. Cortices were dissected out in HBSS, treated for 15 min at 37°C in Trypsin/EDTA-HBSS (Invitrogen), washed once in NeuroBasal medium (NB, Invitrogen) complemented with 10 % Horse Serum to block trypsin activity, washed once more in NB alone. Cortical neurons were dissociated, plated on glass coverslips in NB complemented with B27 complement, 2 mM LGlutamine and 50µg/ml Pencillin/ Streptamycin (InVitrogen) and cultured for 3 days at 37°C, 5% CO2. Coverslips were coated overnight with poly-L-Lysine (10 µg/ml).

15 Transfection and internalisation experiments

[0061] Neurons were electroporated using Amaxa Nucleofactor Kit according to manufacturer's instructions. Hela cells were grown on coverslips and transfected using lipofectamine 2000 (Invitrogen) using manufacturer's protocol. After 8 to 24 hours of transfection cells were processed to study internalisation kinetics or fixed using paraformaldehyde 3%.

[0062] Internalisation assays were performed using FITC conjugated transferrin or unconjugated antibodies. The cells were first incubated for 20 min at 37°C in DMEM/100mM Hepes to eliminate endogenous transferrin. Cells were incubated 15 min at 4°C with ligand and/or antibody and washed twice in icecold PBS before incubation with DMEM, 1 % BSA, 100mM Hepes at 37°C during different times prior fixation and immunocytochemistry. Neurons were processed identically in NB medium.

[0063] Cortical neuron biotinylation was performed using EZ-Link Sulfo-NHS-Biotin kit (PIERCE) with 15 min reaction at 4°C followed by 3 washes with ice-cold PBS containing 10 mM Glycine. Cells were incubated for 5 and 45 min at 37°C to allow endocytosis of biotinylated membrane proteins, prior fixation and immunostaining.

30 Immunoblots and immunoprecipitation

[0064] 1% Triton X-100 cell extracts complemented with protease inhibitors cocktail (Roche) were immunoblotted after separation by 12% or 7-17% gradient SDS-PAGE.

[0065] Immunoprecipitation with anti-flag antibody and N-glycosidase F or endoglycosidase H (Calbiochem) treatment were performed as described (Cappello et al., 2004).

Results

Sequence identification

ESTs coding for overlapping sequences were isolated. After reconstitution of the full length sequence and extensive analysis, the Inventors identified a potential open reading frame (ORF) coding for a new member of the LAMP family (SEQ ID NO: 3). The putative ORF codes for a protein of 280 aa (SEQ ID NO: 4) and is predicted to contain a transmembrane domain (aa 237-256) and a cytoplasmic tail of 24 residues (aa 257-280) (Figure 1). This predicted cytoplasmic domain contains a YKHM motif (aa 276-279) corresponding to a classical YXXO internalization sequence and endosomal targeting signal (Figure 1). The sequence contains 4 highly conserved cysteine residues separated by a fixed number of amino acids which are likely to form the characteristic internal di-sulfide bonds required to form LAMP-like domains. [0067] One of the hallmarks of LAMP proteins is their important level of sugar modification. In agreement, the protein was also predicted to contain 3 consensus N-glycosylation sites. The nucleotide sequence was found to be homologous at the level of 41.6% and 46.4% with LAMP 1 and LAMP 2 respectively, the founding members of the family, while protein alignments displayed 24.7% and 21.2% of identity. Based on its size and the presence of one LAMP domain, the protein was placed on an evolutionary classification tree between LAMP1 and DC-Hil sequences, clearly identifying it as a new member of the LAMP family. DC-Hil (which shares 15.5% identity with LAMP1), is a dendritic cell-specific molecule functioning as an integrin ligand.

[0068] Analysis of the expression of the identified mouse mRNA by northern blot using various mouse tissues indicated that its presence mainly, if not exclusively, in the adult brain (<u>Figure 2</u>). Expression in the embryo was extremely low. The detected mRNA corresponds to a unique transcript of around 1.8kb with no apparent alternative spliced forms.

[0069] Analysis of the expression of the human BAD-LAMP mRNA by northern blot is presented in <u>Figure 3.</u> As in mice, the expression of the mRNA appears to be essentially localized in the brain.

[0070] Based on its relationship to the LAMP family and its restricted pattern of expression, the above-identified protein was named BAD-LAMP, for Brain-AssociateD LAMP.

BAD-LAMP is a glycosylated membrane associated protein

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[0071] To investigate further BAD-LAMP distribution and function, antibodies were raised against peptide epitopes present in its cytoplasmic tail. These antibodies were characterized by immunoblot of HeLa cells transfected with the cDNA coding for mouse BAD-LAMP (Figure 4). Several bands were detected in extracts from transfected cells, while control extracts remained non-reactive.

[0072] Brain extracts displayed also several bands, mostly corresponding to those observed in transfected cells, confirming the existence of several isoforms of BAD-LAMP.

[0073] The detected proteins had a significantly higher molecular weight than the one predicted from the primary sequence of BAD-LAMP (31.7 kDa). In order to define the nature of these post-translational modifications and in absence of immuno-precipitating antibodies, an N-terminally tagged form of BAD-LAMP (FLAG-BAD-LAMP) was transfected allowing efficient immuno-precipitation and treatment with endoglycosidase H (Endo H) and N-glycosidase F (N-gly F). [0074] Immuno-precipitated FLAG-BAD-LAMP was shown to be heavily glycosylated (Figure 5). The major form of the protein (38 kDa, gH) remained Endo H sensitive, thus reflecting endoplasmic reticulum (ER) retention due to over-expression. The two higher additional bands (47 and 53 kDa, gF) were Endo H resistant but remained N-gly F sensitive, as indicated by the accumulation of a fully trimmed 31 kDa protein (g0) after treatment. Transfected BAD-LAMP is therefore heavily glycosylated on at least 2 of its 3 acceptor sites, a situation likely to be shared with endogenous BAD-LAMP detected in brain.

[0075] Although glycosylation was in support of BAD-LAMP membrane association, the membrane bound nature of BAD-LAMP was demonstrated by submitting mouse cortex post-nuclear supernatants to high speed ultracentrifugation in which BAD-LAMP was found associated with the membranes pellets like other membrane associated molecules such as Rab 3a, syntaxin 6 and syntaxin 13 (Figure 6).

[0076] Thus BAD-LAMP, is a glycosylated LAMP-like molecule associated to cortical membranes.

30 BAD-LAMP is expressed in neurons of specific cortical layers after birth.

[0077] Analysis of mouse brain extracts by immunobloting revealed that levels of BAD-LAMP increased strongly after birth (P0) reaching its maximum level at adulthood, but being already strongly expressed at P10-P12 stages (Figure 7). [0078] In situ hybridization was used to investigate in detail the expression pattern of BAD-LAMP in the developing mouse forebrain. First expression of BAD-LAMP was found at P2 in the cingulate cortex, where a thin band of cells in an intermediate position is labelled (Figure 8). At P5, expression extended ventrally into the cortical plate. Furthermore, the caudate putamen showed a punctuate expression of BAD-LAMP transcripts. This expression pattern was maintained at P7, when an additional broad band of large and strongly BAD-LAMP-positive cells appeared in superficial parts of the cortical plate. Although the cortical expression intensified until P9, no major regional changes in BAD-LAMP expression were obvious during this period. At P12 expression of BAD-LAMP in the striatum ceased, while expression further intensified in the cortex. This expression pattern was stable until adulthood.

[0079] Altogether, this expression pattern is in agreement with a function of BAD-LAMP not in early steps of brain development, like neurogenesis and cell migration, but potentially during terminal steps of neuronal differentiation and neuronal function.

[0080] Within the adult cortex, the homogenous staining in outer regions of the cortical plate, as well as in a more restricted band of cells localized centrally was suggestive of an expression in neurons of specific cortical layers. Well-known markers for cortical layers were used to further characterize the respective populations. Comparison of the expression of BAD-LAMP to that of CUX2, a marker for layers II-IV, showed that the BAD-LAMP domain is included in the CUX2 domain and confined to its outer part (Figures 9 and 10). Thus, BAD-LAMP is expressed in the upper layers II and III of the neocortex, but is excluded from layer IV.

[0081] Furthermore, comparison to the layer V marker ER81 (Figures 9 and 10) showed a perfect overlap, demonstrating that the deeply positioned BAD-LAMP-positive population is located in layer V.

[0082] The size of the BAD-LAMP-positive cells in the respective cortical layers was suggestive of neuronal cells. To confirm this observation the expression of BAD-LAMP was investigated in Scrambler mice. These animals show a well described inversion of the layers of cortical projection neurons, with upper layer neurons (layers II-IV) positioned deep while deep layer neurons (layers V and VI) are positioned superficially (Rice and Curran, 1999).

[0083] The organization of BAD-LAMP positive cells in the Scrambler cortex was strikingly altered (Figure 10). Small, lighter stained cell bodies were displaced towards the ventricular side, while larger and stronger labelled cells were

merely found at the pial side of the cortex. This pattern was in agreement with an inversion of the position of BAD-LAMP-positive cells and suggestive of a projection neuron identity. Furthermore, all BAD-LAMP positive cells in the cortex coexpressed the neuronal marker NeuN, again confirming the neuronal identity of labelled cells.

[0084] BAD-LAMP protein distribution in cortical brain sections further confirmed its expression in specific layers. Higher magnification at the confocal level showed that protein was found at the plasma membrane and in vesicles mostly confined to the somatodendritic compartment as delineated by MAP2 staining. Within dendrites, BAD-LAMP was enriched in vesicles clustered in defined domains. BAD-LAMP mostly accumulated within the boundaries of specific neuronal areas *in vivo*.

BAD-LAMP is distributed in specific domains of cortical neurons.

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[0085] In order to confirm the relevance of the above observations, embryonic cortical neurons were explanted and BAD-LAMP sub-cellular distribution was investigated after 3 days of culture. Due the particular clustered distribution of BAD-LAMP, the distribution of proteins known to partition in different cellular domains such as lipid microdomain associated proteins (Madore et al., 1999) was also investigated.

[0086] Using confocal microscopy, it was found that semi-ordered lipid microdomain residents such as PrP, N-CAM, as well as the ganglioside GM-1 (stained with cholera toxin, CT) were enriched in zones excluding BAD-LAMP vesicles. This observation was particularly striking with CT staining and N-CAM, which accumulated almost exclusively in areas negative for BAD-LAMP.

[0087] In contrast, at this low magnification, Thy-1, a molecule representing ordered lipid microdomain-associated proteins, displayed an overlapping distribution with BAD-LAMP.

[0088] However, at higher magnification, no direct co-localization of Thy-1 and BAD-LAMP molecules could be observed. Instead, accumulation of BAD-LAMP-containing vesicles was revealed directly underneath Thy-1 enriched area at the plasma membrane.

[0089] BAD-LAMP containing vesicles therefore accumulate in cellular zones, defined by the presence of Thy-1 at the plasma membrane, while they are segregated away from the detergent resistant microdomains containing most of PrP, GM-1 and N-CAM (Madore et al., 1999).

[0090] BAD-LAMP distribution and microdomain organization seemed to be closely linked. Cholesterol depletion efficiently affects lipid microdomains and was therefore tested for its ability to influence BAD-LAMP distribution.

[0091] Cortical neurons were treated for 2h with cholesterol-oxydase prior to immunostaining and confocal microscopy visualization. As expected, cholesterol depletion had a potent effect on GM-1 distribution at the plasma membrane. Moreover BAD-LAMP vesicular staining was also deeply affected, displaying an extensive co-localization with GM-1, which was never observed in normal conditions. Thus, microdomain organization at the membrane and clustering of BAD-LAMP-positive vesicles appeared to be directly linked. However BAD-LAMP distribution is not dependent on the neuronal polarity.

The cytoskeleton controls the distribution of BAD-LAMP vesicles

[0092] This particular organization was likely to be maintained with the active participation of the cytoskeleton and/or associated proteins. In order to test this hypothesis, several candidate molecules were followed by confocal microscopy in cortical neurons.

[0093] Surprisingly, BAD-LAMP containing-vesicles clustered within punctuate zones delimited by staining with the Smi31 antibody. Smi31 detects phosphorylated epitopes present in neurofilament H and in the microtubule-associated molecule 1 B (Mab1 B) (Fischer and Romano-Clarke, 1990). Although the precise function of MAP1B phosphorylation is still debated, experimental evidences suggest a role in regulating microtubules and actin dynamics as well as being necessary for axonal growth (Dehmelt and Halpain, 2004) (Del Rio et al., 2004). The perfect overlapping distribution of BAD-LAMP and Smi31 strongly suggests that microtubules or actin play an important role in the clustering of BAD-LAMP positive vesicles.

[0094] BAD-LAMP positive vesicles were found in close vicinity of the microtubule network, mirroring by their accumulation the intensity of the tubules. A treatment with the microtubule depolymerizing agent nocodazole was thus carried out. Nocodazole induced a strong redistribution of BAD-LAMP containing vesicles and a loss of BAD-LAMP staining intensity in cortical neurons. Thus the microtubule network influences the positioning of BAD-LAMP vesicles. Lipid microdomain organization and BAD-LAMP distribution in cortical neurons are therefore linked, use the microtubule network and possibly depend on MAP-1 B phosphorylation for their regulation.

BAD-LAMP defines a specific subset of early endosomes.

[0095] The unusual distribution of BAD-LAMP vesicles led the inventors to investigate their relationship with other

types of sub-cellular compartments. The inventors focused primarily on endocytic organelles, likely to be relevant to a transmembrane molecule bearing a YXX Φ motif in its cytoplasmic tail.

[0096] BAD-LAMP could not be detected in classical endosomal compartments (Figure 11), as judged from its lack of co-localization with LAMP-2 (late endosomes and lysosomes), internalized transferrin-FITC or syntaxin 13 (sorting and recycling endosomes (Hirling et al., 2000). BAD-LAMP was not found in more specialized endocytic compartment such as TI-VAMP positive vesicles (Coco et al., 1999). Colabeling with synaptic vesicles proteins such as synaptotagmin I, Rab 3a, VAMP 2 revealed some level of co-localization with BAD-LAMP in the growth cones. Interestingly, co-localization was not observed in other cellular areas and a similar overlapping distribution in the growth cone was observed with TI-VAMP, which is not found enriched in synaptic vesicles (Coco et al., 1999). Thus, this co-distribution in the growth cone probably reflects the difficulty to segregate, at this optical resolution, individual carrier vesicles congregating in the same area of the cone, rather than a true co-localization in the same vesicles. Pre-and post-synaptic transport carriers are derived from trans-Golgi network (TGN) vesicles, which aggregate at initial contacts between axons and dendrites (Sytnyk et al., 2002).

[0097] The inventors therefore attempted to associate BAD-LAMP with other known vesicular markers of these pathways, such as syntaxin 6 or N-CAM (Sytnyk et al., 2004). Co-localization of BAD-LAMP with any of these markers (Figure 11) failed, suggesting that the molecule is sorted in an uncharacterized type of vesicles, which can accumulate in the growth cone of developing axon, as well as in defined and organized domains along the cellular processes.

[0098] BAD-LAMP distribution at the plasma membrane as well as in localized intracellular vesicles suggested a possible shuttling of the molecule between the cell surface and the vesicles. The co-localization of GM-1 and BAD-LAMP upon cholesterol depletion suggests that BAD-LAMP vesicles are accessible to plasma membrane constituents under specific conditions.

[0099] To address this issue, cortical neurons were surface biotinylated at 4°C prior to incubation at 37°C. Biotinylated surface proteins could either diffuse, or be internalized and their intermixing with BAD-LAMP positive compartments was evaluated at different time points by confocal microscopy. Biotinylated proteins were detected rapidly co-localizing with BAD-LAMP after 5 min of internalization. This significant overlapping distribution was decreased after 45 min, suggesting that BAD-LAMP containing organelles could represent a subset of early endocytic vesicles, rapidly accessible from the neuronal surface and serving as an intermediate step for the intracellular sorting of specific surface molecules present in developing neurons.

BAD-LAMP sorting in transfected neurons

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[0100] To further investigate distribution of BAD-LAMP, N- terminally FLAG and C-terminally GFP-tagged BAD-LAMP constructs were generated and their behaviour was monitored by microscopy in co-transfection experiments of cortical neurons

35 [0101] Surprisingly, endogenous BAD-LAMP expression and domain organization were strongly inhibited in electroporated neurons. Nevertheless, transfected FLAG-tagged BAD-LAMP was found enriched in vesicles clustered in specific zones along the neurites.

[0102] Clearly, the tagged-protein is not addressed in conventional endo/lysosomes as judged by its lack of colocalization with LAMP2, internalized transferring or cholera toxin. FLAG-BAD-LAMP exact localization in the cell body was difficult to establish since its over-expression induced an accumulation of the molecule in the ER and Golgi apparatus. Surprisingly the C-terminally GFP-tagged, BAD-LAMP (BAD-GFP) was found accumulating in LAMP2-positive lysosomal compartments. BAD-LAMP cytoplasmic tail contains therefore a cryptic lysosomal retention motif, which is revealed by the addition of the GFP moiety. This observation also suggests that the cytoplasmic tail of BAD-LAMP is actively interacting with, or modified by, molecules which promote its sorting away from traditional endocytic compartments.

[0103] Co-localization of BAD-GFP and FLAG-tagged BAD-LAMP was observed in discrete vesicles in neurites, despite the fact that BAD-GFP was found mostly accumulating in large lysosomes in the cell body. This demonstrates that a small fraction of BAD-GFP can be sorted normally.

[0104] The internalization dynamics of BAD-LAMP was next evaluated by using the N-terminally FLAG-tagged construct and by monitoring FLAG antibody uptake after cold binding. The antibody was rapidly endocytosed after 5 min at 37°C. Inside the cell, it was detected in a compartment different from conventional endo/lysosomes as shown by the absence of co-localization with co-transfected BAD-GFP, LAMP2 and internalized cholera toxin. After 30 min of synchronous uptake, co-localization of the antibodies with BAD-LAMP-GFP and LAMP 2 indicated that BAD-LAMP can reach conventional endocytic compartments, after being internalized from the surface. Surprisingly, this co-localization was more evident in the more discrete LAMP2 positive organelles present in the neurites, than in the large lysosomes observed in the cell body.

[0105] The contribution of tyrosine 276 to BAD-LAMP trafficking was next investigated by introducing a mutational change to alanine at this position (Tyr276Ala). The FLAG-tagged mutant was also found accumulating in the ER and Golgi apparatus of transfected neurons. However, the fraction of the mutant exiting these organelles accumulated at

the surface of the neurites in a manner very distinct from the normal molecule (wt), which was mostly found in intracellular vesicles. Similar results were obtained with a construct lacking the entire cytoplasmic tail of BAD-LAMP. Thus, tyrosine 276 is directly involved in intracellular addressing of BAD-LAMP and allows its internalization from the surface. FLAG antibody uptake after binding in the cold was performed in neurons expressing FLAG-BAD-LAMP Tyr276Ala. Transfected cells remained mostly antibody-decorated at the surface 30 min after warming-up at 37°C. Thus, BAD-LAMP is probably cycling between the plasma membrane and a subset of endocytic vesicles.

BAD-LAMP recycles in HeLa cells.

[0106] In order to further dissect the molecular mechanisms governing BAD-LAMP endocytosis, the distribution and transport of transfected BAD-LAMP was studied in a cell type easy to manipulate, such as HeLa and mouse NIH 3T3 cells. [0107] In HeLa cells, FLAG-BAD-LAMP was found at the cell surface and in internalized transferring containing vesicles distributed in the vicinity of the plasma membrane, while the Tyr276Ala mutant accumulated only at the cell surface (Figure 9A). No co-localization was found in LAMP1-positive late endosomes or lysosomes, nor with co-transfected DC-LAMP tagged with GFP, another lysosomal resident of the LAMP family (de Saint-Vis et al., 1998).

[0108] These observations were confirmed after Percoll density gradient subcellular fractionation of transfected HeLa cells (Figure 12). BAD-LAMP was mostly detected in the low density fractions of the gradient containing plasma membrane, ER and early endosomes, while it was absent from the high density fractions containing lysosomes as indicated by β -hexosaminidase activity.

[0109] Thus, most of transfected BAD-LAMP was found on the cell surface contrasting with transfected neurons in which BAD-LAMP mostly accumulated intracellularly, underlining the specificity of its sorting even when over-expressed. [0110] Anti-FLAG antibody uptake in transfected cells saturated with FITC-transferrin (FITCTF), confirmed that BAD-LAMP could be internalized rapidly in sorting endosomes. Interestingly, 15 min after uptake BAD-LAMP was found present in transferrin positive recycling endosomes clustered around the microtubule organizing center, suggesting that BAD-LAMP could recycle to the plasma membrane after internalization. This hypothesis was supported by the poor colocalization of the antibody with LAMP1 after 45 min of uptake, indicating that the molecule does not reach efficiently late endocytic compartments. This underlines again a difference with neurons, in which the antibodies could be detected in discrete LAMP1 positive compartment 45 min after uptake.

[0111] The molecular mechanisms involved in BAD-LAMP endocytosis were next investigated.

[0112] Experiments performed in cells co-transfected with wild type GTPase dynamin II or dominant negative mutant A44K, indicated that BAD-LAMP internalization is mediated in a dynamin-dependent manner, since antibody internalization was abolished in cells expressing dynamin A44K.

[0113] In order to further define the endocytic pathway used by BAD-LAMP to enter the cell, an RNA inhibition approach was performed to reduce the expression of molecules involved in protein triage from the surface such as the clathrin adaptor AP-2 (Dugast et al., 2005; McCormick et al., 2005). Antibody uptake was monitored by immunostaining and FACS detection after binding at 4°C and internalization at 37°C.

[0114] Cells co-transfected with FLAG-BAD-LAMP and control RNAi plasmid showed rapid internalization of the antibody, while RNAi depletion of AP2 clearly inhibited BAD-LAMP internalization as well as transferrin uptake. In cells depleted for AP2, higher surface levels of BAD-LAMP were also constantly detected, suggesting that BAD-LAMP is internalized constantly through a dynamin/AP2 dependent endocytic pathway.

[0115] Interestingly monitoring of surface anti-flag antibody by FACS also indicated that the molecule was rapidly internalized between 5 and 7.5 min after warming up (Figure 13). Surface levels of antibodies then re-increased after 10min, to be diminished again but with a relatively slower internalization rate. These observations confirm that BAD-LAMP and associated antibodies constantly recycle to the plasma membrane with a relatively high efficiency.

Discussion

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[0116] BAD-LAMP sequence analysis clearly indicates that it represents a new member of the LAMP family. However its expression pattern and intracellular distribution are unconventional compared to other LAMP family members, which show a widespread expression and specifically accumulate in the lysosomes.

[0117] The inventors' observations on BAD-LAMP intracellular distribution are clearly indicative of a strong regulation of its trafficking in a subset of early endosomes.

[0118] The absence of co-localization with transferrin or synaptotagmin 1, as well as late endosomal markers such as LAMP1 suggests that these vesicles represent a distinct class of neuronal endosomes. The kinetics of biotinylated proteins and antibody uptake indicate that they can serve as sorting platforms, prior to transport to other organelles, which are positive for LAMP1, but only represent a minor fraction of the neuronal organelles containing LAMP1.

[0119] The inventors have shown that BAD-LAMP, through an interaction with its YKHM domain, requires dynamin and AP-2 to be internalized and sorted towards the early endocytic recycling pathway of transfected HeLa cells. LAMP1

has also been shown to require the AP-2 adaptor, but its sorting is directed towards lysosomes (Janvier and Bonifacino, 2005). Interestingly, modification of the BAD-LAMP C-terminal domain by GFP affects deeply its transport in neurons and demonstrates the existence of an active sorting pathway in these cells, which normally prevents the accumulation of BAD-LAMP in the lysosomes. The YKHM domain is a relatively weak consensus endosomal/lysosomal addressing signal (Bonifacino and Traub, 2003), although it is also found in CTLA-4, a molecule known to recycle upon activation of T cells (Linsley et al., 1996).

[0120] As suggested by its early endosomes distribution, it could be shown that BAD-LAMP also recycles in transfected HeLa cells. Whether this is the case in neurons remains to be further investigated, although it clearly indicates that the "YKHM" domain is not normally used as a lysosomal addressing signal.

[0121] One of the features of BAD-LAMP containing organelles is their clustered distribution. This distribution mirrors the organization of the different microdomains at the cell surface. Whether BAD-LAMP containing organelles participate in the maintenance of this organization within the neuritic plasma membrane remains to be definitely proven. Nevertheless their sensitivity to cholesterol depleting drugs suggests that microdomains and BAD-LAMP containing vesicles are functionally linked. Strikingly, the clustering of the BAD-LAMP containing vesicles is also defined by the distribution of the phosphorylated epitopes (SMI31) found on the microtubule associated protein MAP1B or Neurofilament H (Fischer and Romano-Clarke, 1990). MAP1 B in the cortex has been strongly implicated in synapse formation and function (Kawakami et al., 2003). Such a role has been functionally demonstrated in mice lacking the phosphorylated form of MAP1B specifically in the hippocampus, which show deficits in long term potentiation in the Schaeffer collaterals pathway (Zervas et al., 2005). It is conceivable that MAP1 B is implicated in the positioning and transport of BADLAMP vesicles at sites of postsynaptic densities on the dendrites of cortical neurons, and that this process could be essential for stabilization, function and plasticity of cortical synapses. This is also supported by the timing of BAD-LAMP expression. [0122] Synaptogenesis in the cortex occurs in the postnatal animal with a peak between P10 and P15 to approach adult values (Micheva and Beaulieu, 1996). This increase in functional synapses in the cortex is strikingly mirrored by the expression of BADLAMP. Thus, it appears possible that BAD-LAMP is involved in the terminal maturation steps of defined cortical neuron populations. The transformation of a transient contact between two neurons into a stable and functional synapse requires major changes in the membrane composition of the respective neuronal surface areas.

[0123] Endocytic processes have been implicated in the regulation of synaptic function and plasticity in vertebrates (Vissel et al., 2001) and in drosophila (Dickman et al., 2006). For example, NMDA receptors are subject to constitutive (Roche et al., 2001) as well as agonist-induced (Vissel et al., 2001) internalization through clathrin mediated endocytosis. Interestingly, *in situ* hybridization for NMDAR1 resulted in strong cellular labeling in neurons of layers II/III, V and VI (Rudolf et al., 1996), resembling the pattern which the inventors found for BAD-LAMP in the postnatal cortex. The BADLAMP containing endocytic compartments could therefore play a regulatory role in these events by maintaining specific zones in the neuronal projections.

35 Example 2

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Anti-BAD-LAMP monoclonal antibody production

1. Antibodies to the cytoplasmic part of BAD-LAMP

[0124] Mice and rats were immunized with a peptide of sequence KMTANQVQIPRDRSQYKHMC (SEQ ID NO: 20) which is present both in mice and human BAD-LAMP.

[0125] Animals were first injected intra-peritonealy (IP) with a mixture of 7.5 μ g (mice) or 50 μ g (rats) of peptide coupled to ovalbumine, by a SPdP or glutaraldehyde-based method (a mix of both), in 500 μ L of water and 500 μ L of complete Freund adjuvant (CFA).

[0126] Three other intra-peritoneal injections were performed every three weeks with identical mixtures, except for incomplete Freund adjuvant (IFA) in replacement of CFA.

[0127] Three weeks after the last intra-peritoneal injection, three daily intra-venous (IV) injections were performed with 20 μ g of peptide in 200 μ L by the caudal vein route (mice) or 50 μ g of peptide in 200 μ L by the penis (rats). The last day, animals were sacrificed to get spleen cells.

2. Antibodies to the extra-cellular part of BAD-LAMP

[0128] A vaccination scheme similar to that used for the generation of antibodies to the cytoplasmic part of human BAD-LAMP was devised except that the peptide was replaced by HeLa 229 transfected with a complete human BAD-LAMP expression system. This construct is characterized by a FLAG tag (Sigma) fused with BAD-LAMP at the N-terminal end. Briefly, 10⁶ (mice) or 10⁷ (rats) 16-hours transfected HeLa 229 cells were used for the IP injections, and 10⁶ (mice and rats) 16-hours transfected HeLa 229 cells were used for the IV injections.

3. Hybridoma generation

- [0129] Spleen cells from immunized rats and mice were fused with X63 myeloma cells Kohler & Milstein (1975) Nature 256:495 according to methods well known in the art.
- [0130] Hybridoma clones yielding antibodies to the cytoplasmic part of BAD-LAMP were screened by ELISA with the Aquabind kit (Iwaki Glass Co. Ltd., Tokyo, Japan) following the manufacturer's instructions. Briefly, the immunoreactivity of conditioned media from the hybridoma clones towards the immunizing peptide was assessed by ELISA. Immunoreactive clones were then selected and went through a second round of cloning and immunoreactive selection. The clones the conditioned medium of which was shown to be immunoreactive were then expanded.
- [0131] 4 anti-BAD-LAMP clones targeting the cytoplasmic domain of mouse and human BAD-LAMP were thus recovered including mBAD 14.2 which has been deposited at the CNCM (Institut Pasteur, Paris, France) under the Budapest Treaty on October 19, 2006 with reference number I-3684.
 - **[0132]** The conditioned media of clones yielding antibodies to the extracellular part of BAD-LAMP were screened by FACS using CHO cells expressing the FLAG-BAD-LAMP produced as described in section 2. Immunoreactive clones were then selected and went through a second round of cloning and immunoreactive selection. The clones the conditioned medium of which was shown to be immunoreactive were then expanded.
 - **[0133]** 1 rat anti-human BAD-LAMP clone targeting the extracellular domain of human BAD-LAMP was thus recovered, hBAD 15F8, which has been deposited at the CNCM (Institut Pasteur, Paris, France) under the Budapest Treaty on October 19, 2006 with reference number I-3679.

References

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- Andrejewski N, Punnonen EL, Guhde G, Tanaka Y, Lullmann-Rauch R, Hartmann D, von Figura K, Saftig P (1999)

 Normal lysosomal morphology and function in LAMP-1-deficient mice. J. Biol. Chem. 274:12692-12701.
 - Bonifacino JS, Traub LM (2003) Signals for sorting of transmembrane proteins to endosomes and lysosomes. Annu Rev Biochem 72:395-447.
 - Cappello F, Gatti E, Camossetto V, David A, Lelouard H, Pierre P (2004) Cystatin F is secreted, but artificial modification of its C-terminus can induce its endocytic targeting. Exp Cell Res 297:607-618.
 - Coco S, Raposo G, Martinez S, Fontaine JJ, Takamori S, Zahraoui A, Jahn R, Matteoli M, Louvard D, Galli T (1999) Subcellular localization of tetanus neurotoxin-insensitive vesicle-associated membrane protein (VAMP)/VAMP7 in neuronal cells: evidence for a novel membrane compartment. J Neurosci 19:9803-9812.
 - de Saint-Vis B, Vincent J, Vandenabeele S, Vanbervliet B, Pin JJ, Ait-Yahia S, Patel S, Mattei MG, Banchereau J, Zurawski S, Davoust J, Caux C, Lebecque S (1998) A novel lysosome-associated membrane glycoprotein, DC-LAMP, induced upon DC maturation, is transiently expressed in MHC class II compartment. Immunity 9:325-336. Dehmelt L, Halpain S (2004) Actin and microtubules in neurite initiation: are MAPs the missing link? J Neurobiol 58: 18-33
 - Del Rio JA, Gonzalez-Billault C, Urena JM, Jimenez EM, Barallobre MJ, Pascual M, Pujadas L, Simo S, La Torre A, Wandosell F, Avila J, Soriano E (2004) MAP1 B is required for Netrin 1 signaling in neuronal migration and axonal guidance. Curr Biol 14:840-850.
 - Dickman DK, Lu Z, Meinertzhagen IA, Schwarz TL (2006) Altered synaptic development and active zone spacing in endocytosis mutants. Curr Biol 16:591-598.
 - Dugast M, Toussaint H, Dousset C, Benaroch P (2005) AP2 clathrin adaptor complex, but not AP1, controls the access of the major histocompatibility complex (MHC) class II to endosomes. J Biol Chem 280:19656-19664.
 - Eskelinen EL, Tanaka Y, Saftig P (2003) At the acidic edge: emerging functions for lysosomal membrane proteins. Trends Cell Biol 13:137-145.
 - Eskelinen EL, Illert AL, Tanaka Y, Schwarzmann G, Blanz J, Von Figura K, Saftig P (2002) Role of LAMP-2 in lysosome biogenesis and autophagy. Mol Biol Cell 13:3355-3368.
- Fischer I, Romano-Clarke G (1990) Changes in microtubule-associated protein MAP1 B phosphorylation during rat brain development. J Neurochem 55:328-333.
 - Gruenberg J, Stenmark H (2004) The biogenesis of multivesicular endosomes. Nat Rev Mol Cell Biol 5:317-323. Hirling H, Steiner P, Chaperon C, Marsault R, Regazzi R, Catsicas S (2000) Syntaxin 13 is a developmentally regulated SNARE involved in neurite outgrowth and endosomal trafficking. Eur J Neurosci 12:1913-1923.
- Janvier K, Bonifacino JS (2005) Role of the endocytic machinery in the sorting of lysosome-associated membrane proteins. Mol Biol Cell 16:4231-4242.
 - Kawakami S, Muramoto K, Ichikawa M, Kuroda Y (2003) Localization of microtubule-associated protein (MAP) 1B in the postsynaptic densities of the rat cerebral cortex. Cell Mol Neurobiol 23:887-894.

- Kennedy MJ, Ehlers MD (2006) Organelles and trafficking machinery for postsynaptic plasticity. Annu Rev Neurosci 29:325-362.
- Lin JH, Saito T, Anderson DJ, Lance-Jones C, Jessell TM, Arber S (1998) Functionally related motor neuron pool and muscle sensory afferent subtypes defined by coordinate ETS gene expression. Cell 95:393-407.
- Linsley PS, Bradshaw J, Greene J, Peach R, Bennett KL, Mittler RS (1996) Intracellular trafficking of CTLA-4 and focal localization towards sites of TCR engagement. Immunity 4:535-543.
 - Madore N, Smith KL, Graham CH, Jen A, Brady K, Hall S, Morris R (1999) Functionally different GPI proteins are organized in different domains on the neuronal surface. Embo J 18:6917-6926.
 - Maxfield FR, Tabas I (2005) Role of cholesterol and lipid organization in disease. Nature 438:612-621.

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- McCormick PJ, Martina JA, Bonifacino JS (2005) Involvement of clathrin and AP-2 in the trafficking of MHC class II molecules to antigen-processing compartments. Proc Natl Acad Sci US A 102:7910-7915.
 - Micheva KD, Beaulieu C (1996) Quantitative aspects of synaptogenesis in the rat barrel field cortex with special reference to GABA circuitry. J Comp Neurol 373:340-354.
 - Rice DS, Curran T (1999) Mutant mice with scrambled brains: understanding the signaling pathways that control cell positioning in the CNS. Genes Dev 13:2758-2773.
 - Roche KW, Standley S, McCallum J, Dune Ly C, Ehlers MD, Wenthold RJ (2001) Molecular determinants of NMDA receptor internalization. Nat Neurosci 4:794-802. Rudolf GD, Cronin CA, Landwehrmeyer GB, Standaert DG, Penney JB, Jr., Young AB (1996) Expression of N-methyl-D-aspartate glutamate receptor subunits in the prefrontal cortex of the rat. Neuroscience 73:417-427.
- Saftig P, Tanaka Y, Lullmann-Rauch R, von Figura K (2001) Disease model: LAMP-2 enlightens Danon disease. Trends Mol Med 7:37-39.
 - Santuccione A, Sytnyk V, Leshchyns'ka I, Schachner M (2005) Prion protein recruits its neuronal receptor NCAM to lipid rafts to activate p59fyn and to enhance neurite outgrowth. J Cell Biol 169:341-354.
- Shikano S, Bonkobara M, Zukas PK, Ariizumi K (2001) Molecular cloning of a dendritic cell-associated transmembrane protein, DC-HIL, that promotes RGDdependent adhesion of endothelial cells through recognition of heparan sulphate proteoglycans. J Biol Chem 276:8125-8134.
 - Sunyach C, Jen A, Deng J, Fitzgerald KT, Frobert Y, Grassi J, McCaffrey MW, Morris R (2003) The mechanism of internalization of glycosylphosphatidylinositolanchored prion protein. Embo J 22:3591-3601.
 - Sytnyk V, Leshchyns'ka I, Dityatev A, Schachner M (2004) Trans-Golgi network delivery of synaptic proteins in synaptogenesis. J Cell Sci 117:381-388.
 - Sytnyk V, Leshchyns'ka I, Delling M, Dityateva G, Dityatev A, Schachner M (2002) Neural cell adhesion molecule promotes accumulation of TGN organelles at sites of neuron-to-neuron contacts. J Cell Biol 159:649-661.
 - Tanaka Y, Guhde G, Suter A, Eskelinen EL, Hartmann D, Lullmann-Rauch R, Janssen PM, Blanz J, von Figura K, Saftig P (2000) Accumulation of autophagic vacuoles and cardiomyopathy in LAMP-2-deficient mice. Nature 406: 902-906.
 - Tiveron MC, Hirsch MR, Brunet JF (1996) The expression pattern of the transcription factor Phox2 delineates synaptic pathways of the autonomic nervous system. J Neurosci 16:7649-7660.
 - Vissel B, Krupp JJ, Heinemann SF, Westbrook GL (2001) A use-dependent tyrosine dephosphorylation of NMDA receptors is independent of ion flux. Nat Neurosci 4:587-596.
- Zervas M, Opitz T, Edelmann W, Wainer B, Kucherlapati R, Stanton PK (2005) Impaired hippocampal long-term potentiation in microtubule-associated protein 1 B-deficient mice. J Neurosci Res 82:83-92.
 - Zimmer C, Tiveron MC, Bodmer R, Cremer H (2004) Dynamics of Cux2 expression suggests that an early pool of SVZ precursors is fated to become upper cortical layer neurons. Cereb Cortex 14:1408-1420.

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Claims

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- 25 1. An ex vivo method for marking neurones or neurone substructures of a mammal, comprising detecting the expression of a protein represented by SEQ ID NO: 4 (Mus musculus BAD-LAMP) or by a homologous sequence which presents at least 60% identity to SEQ ID NO: 4, provided that the homologous sequence represents a protein which can be found associated to plasma and/or vesicular membranes within cortical neurones of said mammal.
- 30 **2.** An ex vivo method according to claim 1, wherein cerebral cortical layers are marked.
 - 3. An *ex vivo* method according to claim 1 or 2, wherein the mammal is a human and the protein is represented by SEQ ID NO: 2.
- 4. An *ex vivo* method according to any of claims 1 to 3, wherein the expression of the protein is detected by a ligand comprising at least one binding moiety, which at least one binding moiety is specific for said protein.
 - 5. An ex vivo method according to claim 4, wherein the at least one binding moiety is specific for a protein represented by a sequence selected from the group consisting of SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, SEQ IDNO: 8 and SEQ ID NO: 10.
 - **6.** An *ex vivo* method according to claim 4 or 5, wherein the at least one binding moiety is selected from the group consisting of an antibody, an antigen-specific antibody fragment, a scFv, or an aptamer.
- 45 7. An ex vivo method according to any of claims 4 to 6, wherein the at least one binding moiety is a monoclonal antibody.
 - **8.** An *ex vivo* method according to claim 7, wherein the monoclonal antibody is produced by one of the following hybridomas:
- a hybridoma deposited at the Collection Nationale de Cultures de Microorganismes (CNCM, Paris, France) under the Budapest Treaty on October 19, 2006 with accession number I-3679, or
 - a hybridoma deposited at the Collection Nationale de Cultures de Microorganismes (CNCM, Paris, France) under the Budapest Treaty on October 19, 2006 with accession number I-3684.
- 9. An ex vivo method according to any of claims 4 to 8, wherein the ligand further comprises at least one detectable marker.
 - 10. An ex vivo method according to claim 9, wherein the at least one detectable marker is selected from the group

EP 1 933 149 A1

consisting of an enzyme, a chromophore, or a fluorophore.

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- **11.** An *ex vivo* method according to claim 9 or 10, wherein the ligand consists of at least one binding moiety linked to at least one detectable marker.
- 12. An ex vivo method according to any of claims 4 to 8, wherein the ligand consists of at least one binding moiety.
- **13.** An ex *vivo* method according to any of claims 1 to 3, wherein the messenger RNA (mRNA) which encodes the above-defined protein, or a precursor thereof, is detected.
- **14.** An *ex vivo* method according to claim 13, wherein the mRNA, or precursor thereof, is detected by a detection nucleic acid, which detection nucleic acid is liable to hybridize with a nucleic acid encoding the protein.
- **15.** An *ex vivo* method according to claim 14, wherein the detection nucleic acid hybridizes under stringent conditions with a nucleic acid encoding the protein.
- **16.** An *ex vivo* method according to claim 13 or 14, wherein the detection nucleic acid is selected from the group constituted of nucleic acids comprising or constituted of the complementary sequences of SEQ ID NO: 1, 3, 5; 7 and 9 or fragments thereof having at least 9 nucleotides.
- **17.** An *ex vivo* method according to claim 13, wherein the mRNA, or precursor thereof, is detected by a nucleic acid amplification method.

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EP 1 933 149 A1

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Figure 1

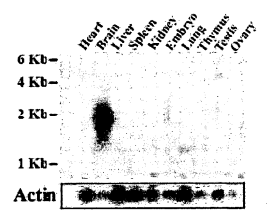


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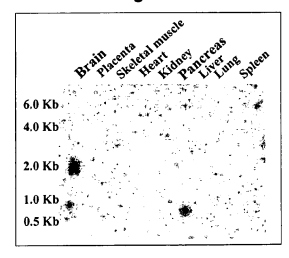


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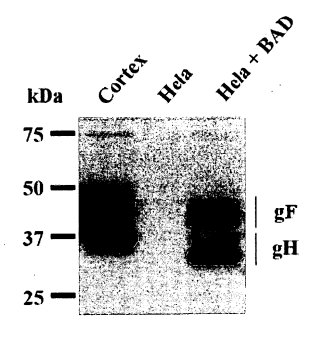


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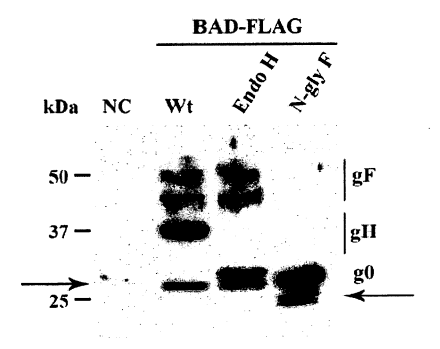


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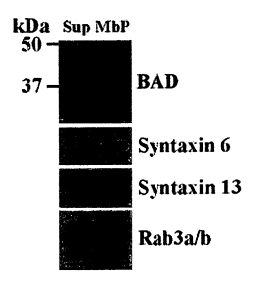


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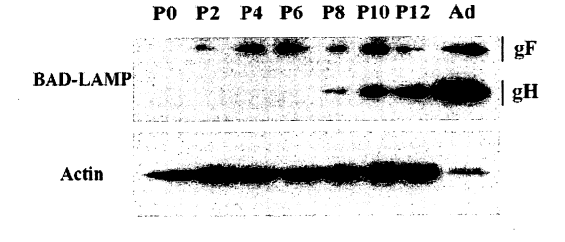
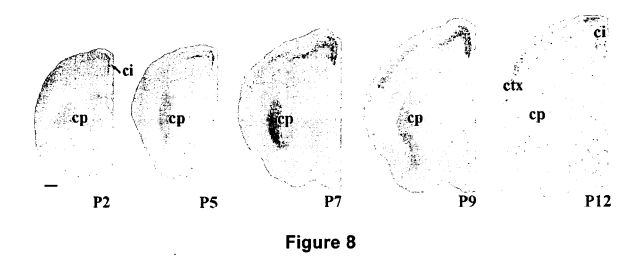


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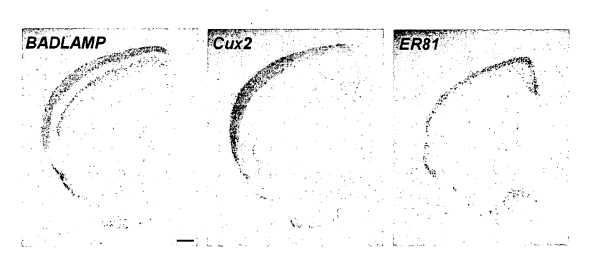


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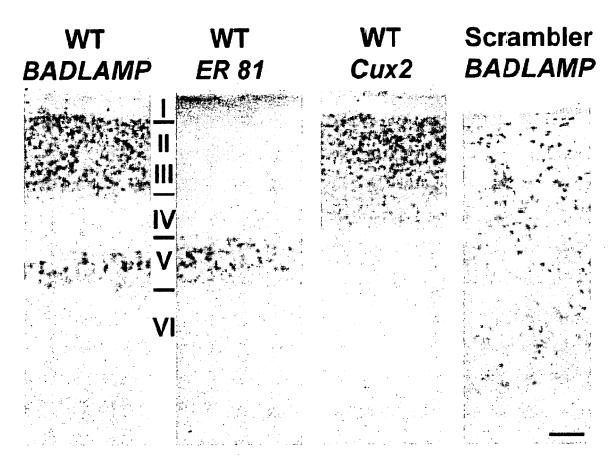


Figure 10

BAD-LAMP/marker

co-localisation Global **Growth cone** Marker Compartment Staining **Staining** Late endosomes LAMP-2 Lysosomes early endosomes TF specific + Ti-VAMP compartment Syt I synaptic vesicles early endosomes Syntaxin 6 **TGN** synaptic vesicles VAMP-2 + synaptic vesicles Rab3 + Recycling Syntaxin 13 endosomes

Figure 11

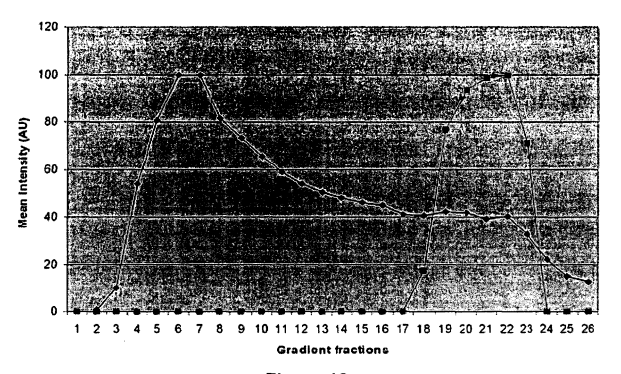


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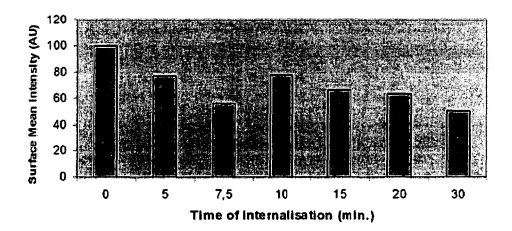


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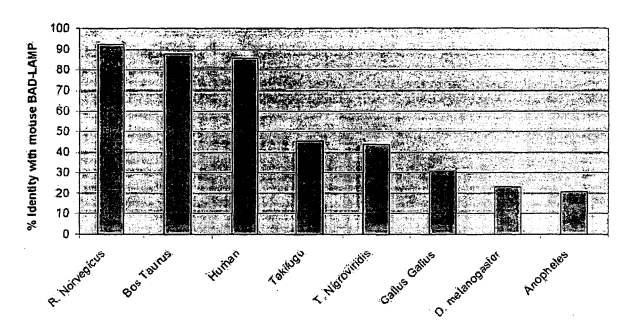


Figure 14



EUROPEAN SEARCH REPORT

Application Number EP 06 29 1947

Category	Citation of document with indication of relevant passages	n, where appropriate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (IPC)	
A	DATABASE EMBL-EBI [Online] XP002426616 retrieved from SRS.EBI.AC.UK/SRSBIN/CGI-BIN/WGETZ?-E+[UNI PROT-ID:CT103 MOUSE] [UNIPROT-ACC:CT103 MOUSE]+-NOSESSION * the whole document *		1-17	INV. G01N33/569	
D,A	ESKELINEN EEVA-LIISA ET acidic edge: Emerging f lysosomal membrane prot TRENDS IN CELL BIOLOGY, vol. 13, no. 3, March 2 pages 137-145, XP002426 ISSN: 0962-8924 * the whole document *	unctions for eins." 003 (2003-03),	1-17		
T	DAVID ALEXANDRE ET AL: a subset of early endoc subpopulations of corti neurons" JOURNAL OF CELL SCIENCE vol. 120, no. 2, Januar pages 353-365, XP009081 ISSN: 0021-9533 * the whole document *	ytic organelles in cal projection , y 2007 (2007-01), 197	1-17	TECHNICAL FIELDS SEARCHED (IPC) G01N	
	Place of search	Date of completion of the search		Examiner	
Munich		23 May 2007	Pellegrini, Paolo		
CATEGORY OF CITED DOCUMENTS X: particularly relevant if taken alone Y: particularly relevant if combined with another document of the same category A: technological background		T : theory or principle E : earlier patent doo after the filing date D : dooument cited in	T: theory or principle underlying the invention E: earlier patent document, but published on, or after the filing date D: document cited in the application L: document cited for other reasons		

REFERENCES CITED IN THE DESCRIPTION

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Non-patent literature cited in the description

- ESKELINEN et al. Trends Cell Biol, 2003, vol. 13, 137 [0005]
- MOLNAR et al. European Journal of Neuroscience, 2006, vol. 23, 921-934 [0016]
- KOHLER; MILSTEIN. Nature, 1975, vol. 256, 495
 [0033] [0129]
- SAMBROOK; RUSELL. Molecular Cloning: A Laboratory Manual. Cold Spring Harbor Laboratory Press, 2001 [0041]
- ANDREJEWSKI N; PUNNONEN EL; GUHDE G; TANAKA Y; LULLMANN-RAUCH R; HARTMANN D; VON FIGURA K; SAFTIG P. Normal lysosomal morphology and function in LAMP-1-deficient mice. J. Biol. Chem., 1999, vol. 274, 12692-12701 [0134]
- BONIFACINO JS; TRAUB LM. Signals for sorting of transmembrane proteins to endosomes and lysosomes. Annu Rev Biochem, 2003, vol. 72, 395-447 [0134]
- CAPPELLO F; GATTI E; CAMOSSETTO V; DAV-ID A; LELOUARD H; PIERRE. Cystatin F is secreted, but artificial modification of its C-terminus can induce its endocytic targeting. Exp Cell Res, 2004, vol. 297, 607-618 [0134]
- COCOS; RAPOSOG; MARTINEZS; FONTAINE
 JJ; TAKAMORIS; ZAHRAOUIA; JAHNR; MATTEOLIM; LOUVARDD; GALLIT. Subcellular localization of tetanus neurotoxin-insensitive vesicle-associated membrane protein (VAMP)/VAMP7 in
 neuronal cells: evidence for a novel membrane compartment. J Neurosci, 1999, vol. 19, 9803-9812
 [0134]
- DE SAINT-VIS B; VINCENT J; VANDENABEELE S; VANBERVLIET B; PIN JJ; AIT-YAHIA S; PATEL S; MATTEI MG; BANCHEREAU J; ZURAW-SKI S. A novel lysosome-associated membrane glycoprotein, DC-LAMP, induced upon DC maturation, is transiently expressed in MHC class II compartment. Immunity, 1998, vol. 9, 325-336 [0134]
- DEHMELT L; HALPAIN S. Actin and microtubules in neurite initiation: are MAPs the missing link. J Neurobiol, 2004, vol. 58, 18-33 [0134]
- DEL RIO JA; GONZALEZ-BILLAULT C; URENA JM; JIMENEZ EM; BARALLOBRE MJ; PAS-CUAL M; PUJADAS L; SIMO S; LA TORRE A; WANDOSELL F. MAP1 B is required for Netrin 1 signaling in neuronal migration and axonal guidance. Curr Biol, 2004, vol. 14, 840-850 [0134]

- DICKMAN DK; LU Z; MEINERTZHAGEN IA;
 SCHWARZ TL. Altered synaptic development and active zone spacing in endocytosis mutants. Curr Biol, 2006, vol. 16, 591-598 [0134]
- DUGAST M; TOUSSAINT H; DOUSSET C; BENAROCH P. AP2 clathrin adaptor complex, but not AP1, controls the access of the major histocompatibility complex (MHC) class II to endosomes. *J Biol Chem*, 2005, vol. 280, 19656-19664 [0134]
- ESKELINEN EL; TANAKA Y; SAFTIG P. At the acidic edge: emerging functions for lysosomal membrane proteins. *Trends Cell Biol*, 2003, vol. 13, 137-145 [0134]
- SCHWARZMANN G; BLANZ J; VON FIGURA K; SAFTIG P. Role of LAMP-2 in lysosome biogenesis and autophagy. *Mol Biol Cell*, 2002, vol. 13, 3355-3368 [0134]
- FISCHER I; ROMANO-CLARKE G. Changes in microtubule-associated protein MAP1 B phosphorylation during rat brain development. *J Neurochem*, 1990, vol. 55, 328-333 [0134]
- GRUENBERG J; STENMARK H. The biogenesis of multivesicular endosomes. Nat Rev Mol Cell Biol, 2004, vol. 5, 317-323 [0134]
- HIRLING H; STEINER P; CHAPERON C; MAR-SAULT R; REGAZZI R; CATSICAS S. Syntaxin 13 is a developmentally regulated SNARE involved in neurite outgrowth and endosomal trafficking. Eur J Neurosci, 2000, vol. 12, 1913-1923 [0134]
- JANVIER K; BONIFACINO JS. Role of the endocytic machinery in the sorting of lysosome-associated membrane proteins. *Mol Biol Cell*, 2005, vol. 16, 4231-4242 [0134]
- KAWAKAMI S; MURAMOTO K; ICHIKAWA M; KURODA Y. Localization of microtubule-associated protein (MAP) 1B in the postsynaptic densities of the rat cerebral cortex. *Cell Mol Neurobiol*, 2003, vol. 23, 887-894 [0134]
- KENNEDY MJ; EHLERS MD. Organelles and trafficking machinery for postsynaptic plasticity. Annu Rev Neurosci, 2006, vol. 29, 325-362 [0134]
- LIN JH; SAITO T; ANDERSON DJ; LANCE-JONES C; JESSELL TM; ARBER S.
 Functionally related motor neuron pool and muscle sensory afferent subtypes defined by coordinate ETS gene expression. Cell, 1998, vol. 95, 393-407 [0134]

- LINSLEY PS; BRADSHAW J; GREENE J; PEACH R; BENNETT KL; MITTLER RS. Intracellular trafficking of CTLA-4 and focal localization towards sites of TCR engagement. *Immunity*, 1996, vol. 4, 535-543 [0134]
- MADORE N; SMITH KL; GRAHAM CH; JEN A; BRADY K; HALL S; MORRIS R. Functionally different GPI proteins are organized in different domains on the neuronal surface. *Embo J*, 1999, vol. 18, 6917-6926 [0134]
- MAXFIELD FR; TABAS I. Role of cholesterol and lipid organization in disease. *Nature*, 2005, vol. 438, 612-621 [0134]
- MCCORMICK PJ; MARTINA JA; BONIFACINO JS. Involvement of clathrin and AP-2 in the trafficking of MHC class II molecules to antigen-processing compartments. *Proc Natl Acad Sci US A*, 2005, vol. 102, 7910-7915 [0134]
- MICHEVA KD; BEAULIEU C. Quantitative aspects of synaptogenesis in the rat barrel field cortex with special reference to GABA circuitry. *J Comp Neurol*, 1996, vol. 373, 340-354 [0134]
- RICE DS; CURRAN T. Mutant mice with scrambled brains: understanding the signaling pathways that control cell positioning in the CNS. Genes Dev, 1999, vol. 13, 2758-2773 [0134]
- ROCHE KW; STANDLEY S; MCCALLUM J; DUNE LY C; EHLERS MD; WENTHOLD RJ. Molecular determinants of NMDA receptor internalization. Nat Neurosci, 2001, vol. 4, 794-802 [0134]
- RUDOLF GD; CRONIN CA; LANDWEHRMEYER GB; STANDAERT DG; PENNEY JB, JR; YOUNG AB. Expression of N-methyl-D-aspartate glutamate receptor subunits in the prefrontal cortex of the rat. Neuroscience, 1996, vol. 73, 417-427 [0134]
- SAFTIG P; TANAKA Y; LULLMANN-RAUCH R; VON FIGURA K. Disease model: LAMP-2 enlightens Danon disease. *Trends Mol Med*, 2001, vol. 7, 37-39 [0134]
- SANTUCCIONE A; SYTNYK V; LESHCHYNS'KA
 I; SCHACHNER M. Prion protein recruits its neuronal receptor NCAM to lipid rafts to activate p59fyn and to enhance neurite outgrowth. J Cell Biol, 2005, vol. 169, 341-354 [0134]

- SHIKANO S; BONKOBARA M; ZUKAS PK; ARIIZUMI K. Molecular cloning of a dendritic cell-associated transmembrane protein, DC-HIL, that promotes RGDdependent adhesion of endothelial cells
 through recognition of heparan sulphate proteoglycans. J Biol Chem, 2001, vol. 276, 8125-8134 [0134]
- SUNYACH C; JEN A; DENG J; FITZGERALD KT; FROBERTY; GRASSIJ; MCCAFFREY MW; MORRIS R. The mechanism of internalization of glycosylphosphatidylinositolanchored prion protein. Embo J, 2003, vol. 22, 3591-3601 [0134]
- SYTNYK V; LESHCHYNS'KA I; DITYATEV A; SCHACHNER M. Trans-Golgi network delivery of synaptic proteins in synaptogenesis. J Cell Sci, 2004, vol. 117, 381-388 [0134]
- SYTNYK V; LESHCHYNS'KA I; DELLING M; DITYATEVA G; DITYATEV A; SCHACHNER M.
 Neural cell adhesion molecule promotes accumulation of TGN organelles at sites of neuron-to-neuron contacts. J Cell Biol, 2002, vol. 159, 649-661 [0134]
- TANAKA Y; GUHDE G; SUTER A; ESKELINEN EL; HARTMANN D; LULLMANN-RAUCH R; JANSSEN PM; BLANZ J; VON FIGURA K; SAFT-IG P. Accumulation of autophagic vacuoles and cardiomyopathy in LAMP-2-deficient mice. Nature, 2000, vol. 406, 902-906 [0134]
- TIVERON MC; HIRSCH MR; BRUNET JF. The expression pattern of the transcription factor Phox2 delineates synaptic pathways of the autonomic nervous system. J Neurosci, 1996, vol. 16, 7649-7660 [0134]
- VISSEL B; KRUPP JJ; HEINEMANN SF; WEST-BROOK GL. A use-dependent tyrosine dephosphorylation of NMDA receptors is independent of ion flux. Nat Neurosci, 2001, vol. 4, 587-596 [0134]
- ZERVAS M; OPITZ T; EDELMANN W; WAINER B; KUCHERLAPATI R; STANTON PK. Impaired hippocampal long-term potentiation in microtubule-associated protein 1 B-deficient mice. J Neurosci Res, 2005, vol. 82, 83-92 [0134]
- ZIMMER C; TIVERON MC; BODMER R; CREMER H. Dynamics of Cux2 expression suggests that an early pool of SVZ precursors is fated to become upper cortical layer neurons. Cereb Cortex, 2004, vol. 14, 1408-1420 [0134]